	TH AND HUMAN SERVICES G ADMINISTRATION
DISTRICT ADDRESS AND PHONE NUMBER	DATE(S) OF INSPECTION
60 Eighth Street NE	03/18/2013 - 04/02/2013*
Atlanta, GA 30309	FEI NUMBER
(404) 253-1161 Fax: (404) 253-1202	3010078541
Industry Information: www.fda.gov/oc/indu	stry
NAME AND TITLE OF INDIVIDUAL TO WHOM REPORT ISSUED	
TO: Austin E. Gore, Owner/Pharmacist in	Charge
FIRM NAME	STREET ADDRESS
Clinical Specialties Compounding Pharmacy	318 Baston Rd, Suite 103
CITY, STATE, 219 CODE, COUNTRY	TYPE ESTABLISHMENT INSPECTED
Augusta, GA 30907	Producer of Sterile Drug Products

This document lists observations made by the FDA representative(s) during the inspection of your facility. They are inspectional observations, and do not represent a final Agency determination regarding your compliance. If you have an objection regarding an observation, or have implemented, or plan to implement, corrective action in response to an observation, you may discuss the objection or action with the FDA representative(s) during the inspection or submit this information to FDA at the address above. If you have any questions, please contact FDA at the phone number and address above.

DURING AN INSPECTION OF YOUR FIRM WE OBSERVED:

OBSERVATION 1

Each batch of drug product required to be free of objectionable microorganisms is not tested through appropriate laboratory testing.

Specifically,

Your firm does not perform sterility and endotoxin testing for any sterile drug products produced at and distributed by your firm. As such, there is no assurance that your aseptic process is effective in achieving sterility of finished, critical drug products. For example:

- 1. Commercially available Avastin 100mg/4-mL vial, lot # (b) (4), was repacked into individual 0.1 mL syringes, lot # CABDBDAC:17, by you on 2/13/13. This lot was released without sterility or endotoxin testing. It is associated with 4 cases of bacterial eye infections after intraocular administration in Athens, GA.
- 2. Commercially available Avastin 100mg/4-mL vial, lot # (b) (4), was repacked into (b) individual 0.1 mL syringes, lot # CABDBDAC:69, by you on 2/13/3. This lot was released without sterility or endotoxin testing. It is associated with 1 case of bacterial eye infection in South Bend, IN.
- 3. Hydroxyprogresterone Caproate (HPC) is produced from non-sterile components at your firm. This drug product is administered to pregnant women at risk of pre-term delivery. No sterility or endotoxin tests have been performed for any finished lots produced and released for records reviewed within the last 6 months.

OBSERVATION 2

Testing and release of drug product for distribution do not include appropriate laboratory determination of satisfactory conformance to the final specifications and identity and strength of each active ingredient prior to release.

Specifically,

Sterile drug products produced at and distributed by your firm have not been assay tested for potency. As such, there is no assurance that these distributed drug products can produce the desired, maximal effect for patients.

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Pharmacy	
CITY STATE ZIP CODE, COUNTRY	TYPE ESTABLISHMENT INSPECTED
Augusta, GA 30907	Producer of Sterile Drug Products

OBSERVATION 3

Aseptic processing areas are deficient regarding the system for monitoring environmental conditions.

Specifically,

- A. Environmental monitoring of the ISO 5 (b) (4) has not been performed adequately and periodically according to an approved written program. For example:
 - SOP No. 4, Environmental Monitoring of the buffer or clean area and antercom area, has not been approved or implemented by you to date. It states that "in addition to viable and nonviable air sampling, the LAFW and/or Barrier Isolators require contact surface test performed at least (b) (4)" Test results are to be recorded on the sterile product maintenance log. There is no documentation that indicates that you have routinely performed these required tests for microbial organisms.
 - 2. There is no documentation maintained at your firm that demonstrates that the ISO 5 (b) (4) has been surface sampled for microbial contamination from the time period covering 2007 March 12, 2013. Certification Reports from contract testing facilities covering this time period only report that air sampling for particle counts has been conducted within the (b) (4)
 - 3. The (b) attached to the (b) (4) of the ISO 5 (b) (4) that contact products during sterile operations (a critical area prone to contamination) have not been monitored for microbial contamination. You do not perform environmental monitoring with each daily production run to demonstrate that microbial limits have not been exceeded after each sterile operation is performed in the (b)
 - 4. (b) (4) Certification Reports provided by contract testing facilities are deficient in that they do not provide enough detail regarding the specific locations in the ISO 5 (b) (4) that were sampled for air quality analysis. Additionally, the reports do not delineate whether viable or non-viable air particles were sampled. Moreover, the tests were not conducted in accordance with an approved procedure by your firm.
 - Airflow smoke pattern tests that were performed by a contract testing facility and documented in (b) (4)
 Certification Reports from 2007 current are deficient in that they do not demonstrate unidirectional airflow under dynamic conditions.
- B. Personnel monitoring is not performed by you after sterile repacking or production operations.
 - 1. There is no determination as to whether bacterial limits have been exceeded during and after sterile operations in the ISO 5(b) (4). Also, the efficacy of your aseptic procedures cannot be determined.
 - There is no determination that you have performed media fills at least semi-annually inside the ISO 5 (b)
 You have not demonstrated that you can perform sterile operations under conditions that closely simulate the most challenging or stressful conditions encountered during repacking or production of sterile products.
 - SOP 7.007.31, Process Simulation Testing, High Risk, has not been approved or implemented by you to date.
 There is no indication that you have performed this procedure for evaluation of your aseptic technique and the
 cleanliness of the equipment used in your sterile production operations.

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SEE REVERSE OF THIS PAGE	Nicole A. Lloyd, Investigator Jawaid Hamid, Investigator	04/02/2013
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TO: Austin E.	Gore, Owner/Pharmacist in	Charge SIREET AUDINESS	
Clinical Specia	alties Compounding	318 Baston Rd, Suite 103	
Pharmacy			
Augusta, GA 30	907	Producer of Sterile Drug Produ	ata
Augusta, GA 30	1307	Floducer of Scerife Drug Flodu	ccs
A. Initial qualific equipment is: 1. The by anot been The programment instrument. 2. The by another instrument instrument. 3. The mading the machine on a temp outage. A the firm to the firm to the positive of the positive of the positive of the control	cation and routine calibration, maintenant performed or documented to assure 4) qualified, maintained, or cleaned accorrammed sterilization cycle (typically 1) indicators or temperature sensing dev Further, the identification of non-steril ted. Moreover, the 16 (4) has not but in achieving uniform distribution of 14 (4) used for lified, maintained, or cleaned according the terquires (5) (4) sterilization. Not be effectiveness of the (6) There is no for the load sizes to ensure uniform distributions because recording chart. There is no be a formal to the temperature fluctuations because the refrigerator is not monitored for temperature. In the temperature is not monitored for temperature. In the cause of the temperature fluctuations are pressure has been working properly a such as Estradiol Cypionate, Progeste produced from non-sterile components. There is no documentation that (b) in a hangs from a metal rod in the ISO 5 oil-based drug solutions (b) (4) written procedure for the use of the cause.	used for the sterilization of aqueous injectading to the operation's manual or an approved (a) has not been valid ices are used to verify the effectiveness of this e components that enter the machine or the load een temperature mapped to demonstrate the catemperature. (b) (4) sterilization of aqueous injectable solid to an approved written program. Moreover, the color of the non-sterile components of the non-sterile compo	and electronic able solutions has written program. ated. No sterilization d sizes are not pability of the autions has not the temperature reach drug vices are used to that enter the asse of a power anel present at the mand the no assurance that soms. The production run, e to syringes in and vials. There is alking gun is
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FORM FDA 483 (69/08)	PREVIOUS EDITION OBSOLETE INSPE	CTIONAL OBSERVATIONS	PAGE 3 OF 7 PAGES

	TH AND HUMAN SERVICES G ADMINISTRATION
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60 Eighth Street NE	03/18/2013 - 04/02/2013*
Atlanta, GA 30309 (404) 253-1161 Fax:(404) 253-1202	3010078541
Industry Information: www.fda.gov/oc/indu	stry
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TO: Austin E. Gore, Owner/Pharmacist in	STRLET ADDRESS
Clinical Specialties Compounding	318 Baston Rd, Suite 103
Pharmacy CITY, STATE, ZIP CODE, COUNTRY	Type ESTABLISHMENT INSPECTED
Augusta, GA 30907	Producer of Sterile Drug Products
OBSERVATION 5 The flow of components, drug product containers, in-process to prevent contamination. Specifically,	naterials, and drug products though the building is not designed
A. Avastin	
1. Commercially available and sterile vials of Ava time temperature monitoring. Without a backup Avastin products have not been subjected to abn (i.e. power outages over the weekend when perss 2. Materials and supplies such as pre-packaged ster from the ISO 7 buffer area/gowning room and tr mock demonstration of avastin repacking/unit de supplies prior to their entry into the ante chambee chamber of the (b) (4) Only the top portion the mock demonstration) was disinfected with Additionally, approved and implemented SOP 8. Avastin Syringes, does not include the provision product production prior to ante chamber entry of B. Hydroxyprogesterone Caproate (HPC) 1. Non-sterile components used in the production of weighed and mixed in a (b) (4) transfer to the ISO 7 buffer area/gowning room. components used for the production of sterile pre 2. You stated that hold times for aqueous, non-sterile. However, you do not have batch data to substanstiate these hold times for aqueous mixture of HPC non-sterile components ISO 7 buffer area. There is no justification for the that non-sterile, aqueous mixtures are exposed to the buff solutions contained in beakers are further the beaker prior to (b) (4) ante chamber entry C. Medroxyprogesterone 1. Non-sterile components used in the production of the prod	for disinfecting materials and supplies pivotal for sterile drug for disinfecting materials and supplies pivotal for sterile drug for the form of the
SEE REVERSE Nicole A. Lloyd, Investigate Jawaid Hamid, Investigator	04/02/2013
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		LTH AND HUMAN SERVICES	
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	GA 30309 3-1161 Fax:(404) 253-1202	3010078541	
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	tin E. Gore, Owner/Pharmacist in		
FIRM NAME		STREET ADDRESS	
	Specialties Compounding	318 Baston Rd, Suite 103	
Pharmacy city, state, zie co	OF COUNTRY	TYPE ESTABLISHMENT INSPECTED	
Augusta,	GA 30907	Producer of Sterile Drug P	roducts
3.	The aqueous mixture of Medroxyprogesterone located in the ISO 7 buffer area. There is no ju		
OBSERVA	TION 6		
Aseptic proc conditions.	essing areas are deficient regarding the system f	or cleaning and disinfecting the room to p	roduce aseptic
Specifically,			
A. ISO	0.5 (b) (4)		
	There are no written, approved procedures for l	(SO 5 (b) (4) cleaning and frequency as after sterile operations are performed.	nd there is no
2.	You stated that (b) (4) is does not include the ante chamber of the (b) (4) placed for aseptic operations.	where drug products, materials, and s	
3.	You stated that you use commercially available surfaces. The active ingredients for the wipes a These wipes' effectiveness in removing spores from the	actives are not sporicidal and you have no	
B. ISO	7 IV Room and Gowning Room		
	SOP 5.001, Cleaning and Disinfection, states th program of cleaning and disinfection" and that ceilings, floors, all equipment, and working sur by you to date.	"surfaces must be cleaned and disinfected"	to include walls,
	The only documentation of ISO 7 IV room clear February 2012. These logs are deficient in that of cleaning, the type of cleaning solutions used,	they do not include information such as th	ne actual time and date
3.	There is no documentation that the ISO 7 gown	ing room floor has been cleaned/sanitized	on a routine basis.
	The shelf located in the gowning room that con operations is disorderly. Used gowns were observed		e production
There is no d	nclassified Room/Work Bench Area ocumentation that the (b) (4) has been cleaned with a 10% bleach and water so	used for weighing and mixing olution or an acrylic cleaner in accordance	
	EMPLOYECES SIGNATURE	274	DATE ISSUED
EE DEVICE	LaReese K. Thomas, Investig		
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CITY, STATE, ZIP CODE, COUNTRY	TYPE ESTABLISHMENT INSPECTED
Augusta, GA 30907	Producer of Sterile Drug Products
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OBSERVATION 7

There is no written testing program designed to assess the stability characteristics of drug products.

Specifically,.

There is no approved or implemented stability program procedures for Beyond Use Dates (BUDs) assigned to sterile drug products. There is also no stability data to support current BUDs of 60, 90, or 180 days.

OBSERVATION 8

Complaint procedures are deficient in that written complaint records are not maintained in a file designated for drug product complaints.

Specifically,

- 1. There are no written, approved procedures for handling written or oral drug complaints.
- 2. Five (5) cases of bacterial eye infections are associated with Avastin lots repacked on 2/13/13. There is no documentation that you conducted an investigation to identify the root cause of the infections.

OBSERVATION 9

The number of qualified personnel is inadequate to perform the manufacture and processing of each drug product.

Specifically,

There is a (b) ratio of Pharmacist in Charge to Pharmacy Technician at the firm and (b) (4) performs sterile drug production at the firm. There is no second person verification of your gowning routine in the buffer area or your aseptic technique during the production of sterile drug products.

	SEE REVERSE	Nicole A. Lloyd, Inv		04/02/2013
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60 Eighth Street NE	03/18/2013 - 04/02/2013*
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Clinical Specialties Compounding	318 Baston Rd, Suite 103
Pharmacy	
CITY, STATE, 3P CODE, COUNTRY	TYPE ESTABLISHMENT INSPECTED
Augusta, GA 30907	Producer of Sterile Drug Products

OBSERVATION 10

GMP training is not conducted on a continuing basis and with sufficient frequency to assure that employees remain familiar with CGMP requirements applicable to them.

Specifically,

There are no training records at your firm to document that you and the pharmacy technician are current on aseptic techniques and practices for the production of both sterile and non-sterile drug products.

OBSERVATION 11

A system by which the distribution of each lot of drug product can be readily determined to facilitate its recall if necessary, has not been established.

Specifically,

There are no written, approved procedures in place to facilitate a recall of drug products at your firm. During the inspection, a recall of all sterile products at your firm was requested. Since no recall system was in place, there was a delay in contacting and notifying clinics to make patients aware of potential drug product contamination.

* DATES OF INSPECTION:

03/18/2013(Mon), 03/19/2013(Tue), 03/20/2013(Wed), 03/21/2013(Thu), 03/22/2013(Fri), 03/25/2013(Mon), 03/26/2013(Tue), 03/27/2013(Wed), 03/28/2013(Thu), 04/02/2013(Tue)

SEE REVERSE	LaReese K. Thomas, Investigator Nicole A. Lloyd, Investigator Jawaid Hamid, Investigator	04/02/2013
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The observations of objectionable conditions and practices listed on the front of this form are reported:

- 1. Pursuant to Section 704(b) of the Federal Food, Drug and Cosmetic Act, or
- 2. To assist firms inspected in complying with the Acts and regulations enforced by the Food and Drug Administration

Section 704(b) of the Federal Food, Drug, and Cosmetic Act (21 USC 374(b)) provides:

"Upon completion of any such inspection of a factory, warehouse, consulting laboratory, or other establishment, and prior to leaving the premises, the officer or employee making the inspection shall give to the owner, operator, or agent in charge a report in writing setting forth any conditions or practices observed by him which, in his judgement, indicate that any food, drug, device, or cosmetic in such establishment (1) consists in whole or in part of any filthy, putrid, or decomposed substance, or (2) has been prepared, packed, or held under insanitary conditions whereby it may have become contaminated with filth, or whereby it may have been rendered injurious to health. A copy of such report shall be sent promptly to the Secretary."